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NEWS 1	Web Page for STN Seminar Schedule - N. America
NEWS 2	AUG 10 Time limit for inactive STN sessions doubles to 40 minutes
NEWS 3	AUG 18 COMPENDEX indexing changed for the Corporate Source (CS) field
NEWS 4	AUG 24 ENCOMPPLIT/ENCOMPPLIT2 reloaded and enhanced
NEWS 5	AUG 24 CA/CAplus enhanced with legal status information for U.S. patents
NEWS 6	SEP 09 50 Millionth Unique Chemical Substance Recorded in CAS REGISTRY
NEWS 7	SEP 11 WPIDS, WPINDEX, and WPIX now include Japanese FTERM thesaurus
NEWS 8	OCT 21 Derwent World Patents Index Coverage of Indian and Taiwanese Content Expanded
NEWS 9	OCT 21 Derwent World Patents Index enhanced with human translated claims for Chinese Applications and Utility Models
NEWS 10	OCT 27 Free display of legal status information in CA/CAplus, USPATFULL, and USPAT2 in the month of November.
NEWS 11	NOV 23 Addition of SCAN format to selected STN databases
NEWS 12	NOV 23 Annual Reload of IEL Databases

NEWS EXPRESS MAY 26 09 CURRENT WINDOWS VERSION IS V8.4,
AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2009.

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NEWS LOGIN Welcome Banner and News Items

Enter NEWS followed by the item number or name to see news on that specific topic.

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SINCE FILE TOTAL
ENTRY SESSION
0.22 0.22

FILE 'REGISTRY' ENTERED AT 13:59:00 ON 30 NOV 2009
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STRUCTURE FILE UPDATES: 29 NOV 2009 HIGHEST RN 1194232-87-5
DICTIONARY FILE UPDATES: 29 NOV 2009 HIGHEST RN 1194232-87-5

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TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

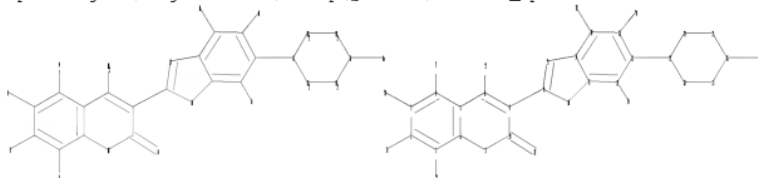
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REGISTRY includes numerically searchable data for experimental and
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on property searching in REGISTRY, refer to:

<http://www.cas.org/support/stngen/stndoc/properties.html>

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Uploading C:\Program Files\Stnexp\Queries\10706328_updated.str



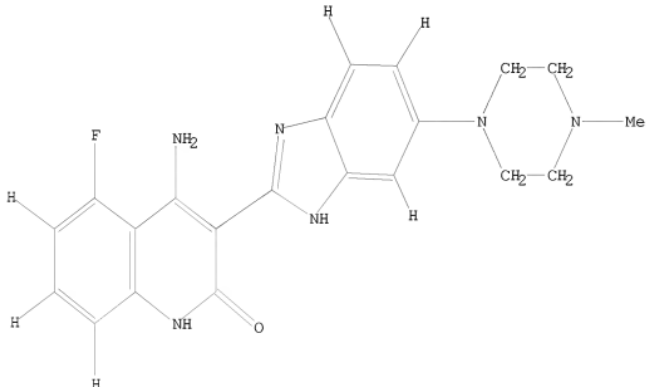
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ring nodes :
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ring bonds :
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16-17 16-19 17-18 17-22 19-20 20-21 21-22 23-24 23-28 24-25 25-26 26-27
27-28
exact/norm bonds :
5-8 6-11 8-9 8-13 9-10 10-11 10-12 14-15 14-18 15-16 17-18 21-23 23-24
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Match level :
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:Atom 9:Atom 10:Atom
11:Atom 12:CLASS 13:CLASS 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom
20:Atom 21:Atom 22:Atom 23:Atom 24:Atom 25:Atom 26:Atom 27:Atom 28:Atom
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L1 STRUCTURE UPLOADED

=> d 11
L1 HAS NO ANSWERS
L1 STR



Structure attributes must be viewed using STN Express query preparation.

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SAMPLE SCREEN SEARCH COMPLETED - 75 TO ITERATE

100.0% PROCESSED 75 ITERATIONS 1 ANSWERS
SEARCH TIME: 00.00.03

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 981 TO 2019
PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

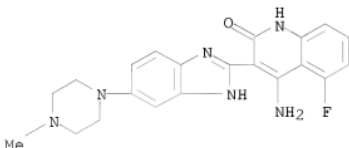
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ED Entered STN: 18 Dec 2006
CN Propanoic acid, 2-hydroxy-, compd. with
4-amino-5-fluoro-3-[6-(4-methyl-1-piperazinyl)-1H-benzimidazol-2-yl]-2(1H)-
quinolinone, hydrate (1:1:1) (CA INDEX NAME)
MF C21 H21 F N6 O . C3 H6 O3 . H2 O
SR CA
LC STN Files: CA, CAPLUS, CHEMCATS, IMSRESEARCH, PHAR, PROUSDDR, SYNTHLINE,
TOXCENTER, USAN

CM 1

CRN 405169-16-6

CMF C21 H21 F N6 O

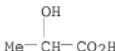


Me

CM 2

CRN 50-21-5

CMF C3 H6 O3



2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> s 11 full

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FULL SCREEN SEARCH COMPLETED - 1493 TO ITERATE

100.0% PROCESSED 1493 ITERATIONS
SEARCH TIME: 00.00.01

34 ANSWERS

L3 34 SEA SSS FUL L1

=> s 11 sss
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SAMPLE SCREEN SEARCH COMPLETED - 75 TO ITERATE

100.0% PROCESSED 75 ITERATIONS
SEARCH TIME: 00.00.01

1 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
PROJECTED ITERATIONS: 981 TO 2019
PROJECTED ANSWERS: 1 TO 80

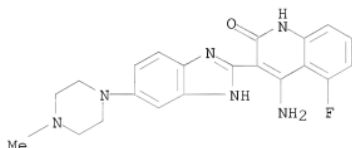
L4 1 SEA SSS SAM L1

=> d 14

L4 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2009 ACS on STN
RN 915769-50-5 REGISTRY
ED Entered STN: 18 Dec 2006
CN Propanoic acid, 2-hydroxy-, compd. with
4-amino-5-fluoro-3-[6-(4-methyl-1-piperazinyl)-1H-benzimidazol-2-yl]-2(1H)-
quinolinone, hydrate (1:1:1) (CA INDEX NAME)
MF C21 H21 F N6 O . C3 H6 O3 . H2 O
SR CA
LC STN Files: CA, CAPLUS, CHEMCATS, IMSRESEARCH, PHAR, PROUSDDR, SYNTHLINE,
TOXCENTER, USAN

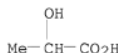
CM 1

CRN 405169-16-6
CMF C21 H21 F N6 O



CM 2

CRN 50-21-5
CMF C3 H6 O3



2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> file caplus
COST IN U.S. DOLLARS SINCE FILE TOTAL
FULL ESTIMATED COST ENTRY SESSION
190.46 190.68

FILE 'CAPLUS' ENTERED AT 14:00:29 ON 30 NOV 2009
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FILE COVERS 1907 - 30 Nov 2009 VOL 151 ISS 23

FILE LAST UPDATED: 29 Nov 2009 (20091129/ED)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

Cplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

CAS Information Use Policies apply and are available at:

<http://www.cas.org/legal/infopolicy.html>

This file contains CAS Registry Numbers for easy and accurate substance identification.

During November, try the new LSUS format of legal status information in the CA/Cplus family databases for free! Complete details on the number of free displays and other databases participating in this offer appear in NEWS 10.

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L5      70 L3

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        758861 ?TUMOR?
        6753 ?TUMOUR?
        6753 ?TUMOUR?
        759258 ?TUMOR?
                  (?TUMOR? OR ?TUMOUR?)
        6753 ?TUMOUR?
        758861 ?TUMOR?
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        759258 ?TUMOUR?
                  (?TUMOUR? OR ?TUMOR?)
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L6      54 L5 AND (?CANCER? OR ?TUMOR? OR ?TUMOUR? OR ?NEOPLASM?)

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L7 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2005:1242789 CAPLUS

DOCUMENT NUMBER: 143:477969

TITLE: Preparation of benzimidazole quinolinones for inhibiting FGFR3 and treating multiple myeloma

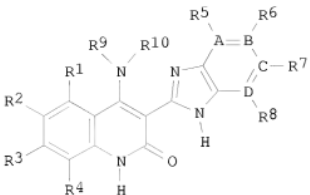
INVENTOR(S): Cai, Shaopei; Chou, Joyce; Harwood, Eric; Heise, Carla C.; Machajewski, Timothy D.; Ryckman, David; Shang, Xiao; Wiesmann, Marion; Zhu, Shuguang
 PATENT ASSIGNEE(S): Chiron Corporation, USA
 SOURCE: U.S. Pat. Appl. Publ., 239 pp., Cont.-in-part of U.S. Ser. No. 644,055.
 CODEN: USXRCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 7
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20050261307	A1	20051124	US 2004-983174	20041105
US 20040092535	A1	20040513	US 2003-644055	20030819 <--
US 7470709	B2	20081230		
CN 1692112	A	20051102	CN 2003-824565	20030819 <--
US 20050203101	A1	20050915	US 2004-839793	20040505
ZA 2006003598	A	20080430	ZA 2006-3598	20060505
US 20090281100	A1	20091112	US 2008-317493	20081223
US 20090181979	A1	20090716	US 2009-398130	20090304
PRIORITY APPLN. INFO.:			US 2002-405729P	P 20020823
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			US 2003-484048P	P 20030701
			US 2003-644055	A2 20030819
			US 2003-517915P	P 20031107
			US 2003-526425P	P 20031202
			US 2003-526426P	P 20031202
			US 2004-546017P	P 20040219
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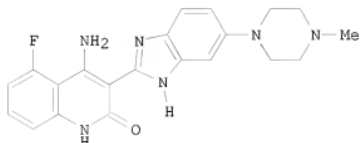
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 143:477969

GI



I



II

AB The title compds. I [A, B, C, and D = C, N; R1-R3 = H, halo, CN, NO₂, etc.; R4 = H, alkyl; R5-R8 = H, halo, CN, NO₂, etc.; R9 = H, (un)substituted alkyl, aryl, etc.; R10 = H], useful for inhibiting fibroblast growth factor receptor 3 or treating a biol. condition mediated by fibroblast growth factor receptor 3, were prepared. E.g., a multi-step synthesis of 4-amino-5-fluoro-3-[6-(4-methylpiperazin-1-yl)-1H-benzimidazol-2-yl]-1H-quinolin-2-one (II), starting from 5-chloro-2-nitroaniline and 1-methylpiperazine, was given. The majority of the exemplary compds. I displayed an IC₅₀ of less than 10 μM with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, Cdk4, MEK1, NEK-2, CHK2, CK1ε, Raf, Fyn, Lck, Rsk2, PAR-1, c-Kit, c-ABL, p60src, FGFR3, FLT-3, PDGFR α , and PDGFR β . In addition, many of the exemplary compds. exhibited IC₅₀ values in the nM range and show potent activity with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, FGFR3, c-Kit, c-ABL, FLT-3, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, MEK1, CHK2, Fyn, Lck, Rsk2, PAR-1, PDGFR α , and PDGFR β with IC₅₀ values of less than 1 μM. The mentioned above compound II was tested in various tests and showed significant antiproliferative activity. II inhibited FGFR3 receptor phosphorylation and ERK phosphorylation in multiple myeloma cell lines with activating FGFR3 mutations.

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

L7 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 20051223876 CAPLUS

DOCUMENT NUMBER: 143:477966

TITLE: Preparation of benzimidazole quinolinones for inhibiting a checkpoint kinase 1 and their use in combination therapy for cancer

INVENTOR(S): Gesner, Thomas G.; Barsanti, Paul A.; Harrison, Stephen D.; Ni, Zhi-Jie; Brammeier, Nathan M.; Zhou, Yasheen; Le, Vincent P.

PATENT ASSIGNEE(S): Chiron Corporation, USA
SOURCE: U.S. Pat. Appl. Publ., 249 pp., Cont.-in-part of U.S. Ser. No. 644,055.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

7

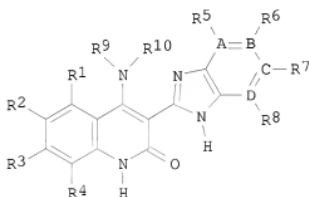
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20050256157	A1	20051117	US 2005-41191	20050121
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US 20090281100	A1	20091112	US 2008-317493	20081223
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		US 2002-428210P	P	20021121
		US 2003-460327P	P	20030403
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		US 2003-460493P	P	20030403
		US 2003-478916P	P	20030616
		US 2003-484048P	P	20030701
		US 2003-644055	A2	20030819
		US 2004-538984P	P	20040123

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): CASREACT 143:477966; MARPAT 143:477966

GI



AB The title compds. [I; A, B, C, D = C, N; R1 = H, halo, CN, NO2, etc.; R2, R3 = H, halo, NO2, CN, etc.; R4 = H, (un)substituted alkyl; R5, R8 = H, (un)substituted alkyl, alkenyl, heterocycl; or R5 may be absent if A = N; or R8 may be absent if D = N; R6, R7 = H, halo, NO2, CN, etc.; R9 = H, (un)substituted alkyl, aryl, etc.; R10 = H; or R9 and R10 join together to form one or more rings, each having 5-7 members], useful for inhibiting checkpoint kinase 1, inducing cell cycle progression, and increasing apoptosis in cells, were prepared. E.g., a multi-step synthesis of 4-amino-3-(benzimidazol-2-yl)-6-(4-methylpiperazinyl)hydroquinolin-2-one, was given. The compds. I were tested against various kinases. Two of the prepared compds. I, 4-[(3S)-1-azabicyclo[2.2.2]oct-3-ylamino]-3-(1H-benzimidazol-2-yl)-6-chloroquinolin-2-(1H)-one and 6-chloro-3-[5-(4-methylpiperazin-1-yl)-1H-benzimidazol-2-yl]-4-[(piperidin-2-ylmethyl)amino]quinolin-2(1H)-one, were found to be potent inhibitors of CHK1 with IC50 of 0.32 nM and 0.63 nM, resp. The majority of the exemplary compds. I displayed an IC50 of less than 10 μ M with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, Cdk4, MEK1, NEK-2, CHK2, CK1 α , Raf, Fyn, Lck, Rsk2, PAR-1, c-Kit, c-ABL,

p60src, FGFR3, FLT-3, PDGFR α , and PDGFR β . In addition, many of the exemplary compds. exhibited IC50 values in the nM range and show potent activity with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, FGFR3, c-Kit, c-ABL, FLT-3, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, MEK1, CHK2, Fyn, Lck, Rsk2, PAR-1, PDGFR α , and PDGFR β with IC50 values of less than 1 μ M. The compds. I may be used to prepare pharmaceutical compns. and may be used in conjunction with DNA damaging agents.

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

L7 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:182836 CAPLUS
DOCUMENT NUMBER: 140:235711

TITLE: Preparation of benzimidazole quinolinolines for inhibiting a serine/threonine kinase
INVENTOR(S): Barsanti, Paul A.; Bussiere, Dirkken; Harrison, Stephen D.; Heise, Carla C.; Jansen, Johanna M.; Jazan, Elisa; Machajewski, Timothy D.; McBride, Christopher; McCrea, William R.; Ng, Simon; Ni, Zhi-Jie; Pecci, Sabina; Pfister, Keith; Ramurthy, Savithri; Renhove, Paul A.; Shafer, Cynthia M.; Silver, Joel B.; Wagman, Allan; Weismann, Marion Chiron Corporation, USA

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 570 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 7

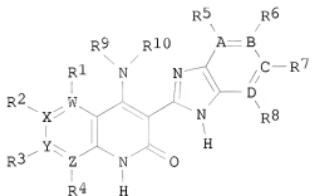
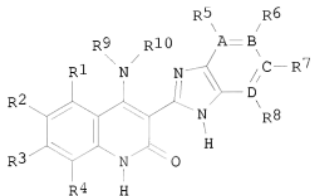
PATENT INFORMATION:

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WO 2004018419	A2	20040304	WO 2003-US25990	20030819 <--
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			US 2003-460493P	P 20030403

US 2003-478916P P 20030616
US 2003-480408P P 20030701
WO 2003-US25990 W 20030819

OTHER SOURCE(S) :
GI

MARPAT 140:235711



AB The title compds. [I and II; A, B, C, and D = C, N; W, X, Y and Z = C, N and at least one of W, X, Y, and Z = N; R1-R8 = H, halo, CN, NO₂, etc.; R9 = H, (un)substituted alkyl, aryl, etc.; R10 = H; or NR9R10 = 5-7 membered ring], useful for inhibiting various enzymes and treating various conditions, were prepared E.g., a multi-step synthesis of 4-amino-3-(benzimidazol-2-yl)-6-(4-methylpiperazinyl)hydroquinolin-2-one, was given. The majority of the exemplary compds. I displayed an IC₅₀ of less than 10 μ M with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, Cdk4, MEK1, NEK-2, CHK2, CK1 α , Raf, Fyn, Lck, Rsk2, PAR-1, c-Kit, c-ABL, p60src, FGFR3, FLT-3, PDGFR α , and PDGFR β . In addition, many of the exemplary compds. exhibited IC₅₀ values in the nM range and show potent activity with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, FGFR3, c-Kit, c-ABL, FLT-3, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, MEK1, CHK2, Fyn, Lck, Rsk2, PAR-1, PDGFR α , and PDGFR β with IC₅₀ values of less than 1 μ M.

OS.CITING REF COUNT: 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L7 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2003:98039 CAPLUS

DOCUMENT NUMBER: 138:153534

TITLE: Preparation of benzimidazolyl-substituted quinolinone derivatives and analogs, with inhibitory action against vascular endothelial growth factor receptor tyrosine kinase, and useful as anticancer agents

INVENTOR(S): Renhowe, Paul A.; Pecchi, Sabina; Machajewski, Timothy D.; Shafer, Cynthia M.; Taylor, Clarke; McCrea,

PATENT ASSIGNEE(S): William R.; McBride, Christopher; Jazan, Elisa
 Chiron Corporation, USA
 SOURCE: U.S. Pat. Appl. Publ., 69 pp., Cont.-in-part of U.S.
 Pat. Appl. 2002 107,392.
 CODEN: USXXCO

DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 2
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20030028018	A1	20030206	US 2002-116117	20020405 <--
US 20020107392	A1	20020808	US 2001-951265	20010911 <--
US 6605617	B2	20030812		
EP 1650203	A1	20060426	EP 2005-17665	20010911 <--
EP 1650203	B1	20080220		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR				
EP 1849782	A1	20071031	EP 2007-11978	20010911 <--
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US 20030158224	A1	20030821	US 2002-284017	20021030 <--
US 6774237	B2	20040810		
US 20040006101	A1	20040108	US 2003-387355	20030312 <--
US 6762194	B2	20040713		
CA 2481055	A1	20031023	CA 2003-2481055	20030404 <--
WO 2003087095	A1	20031023	WO 2003-US10463	20030404 <--
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KE, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
AU 2003226275	A1	20031027	AU 2003-226275	20030404 <--
AU 2003226275	B2	20090723		
EP 1497287	A1	20050119	EP 2003-746614	20030404 <--
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				
BR 2003008996	A	20050222	BR 2003-8996	20030404 <--
CN 1659165	A	20050824	CN 2003-812909	20030404 <--
JP 2005527587	T	20050915	JP 2003-584051	20030404 <--
NZ 536068	A	20080430	NZ 2003-536068	20030404 <--
SG 143985	A1	20080729	SG 2006-6999	20030404 <--
US 20040097545	A1	20040520	US 2003-613411	20030703 <--
US 6800760	B2	20041005		
US 20050054672	A1	20050310	US 2004-886950	20040708
US 7598268	B2	20091006		
MX 2004009739	A	20050111	MX 2004-9739	20041005
IN 2004KN01494	A	20070601	IN 2004-KN1494	20041006
NO 2004004776	A	20041207	NO 2004-4776	20041103
US 20050209456	A1	20050922	US 2005-92137	20050329
US 7335774	B2	20080226		
JP 2007191486	A	20070802	JP 2007-62683	20070312
US 20080070906	A1	20080320	US 2007-866296	20071002
IN 2008KN03126	A	20090206	IN 2008-KN3126	20080730
PRIORITY APPLN. INFO.:			US 2000-232159P	P 20000911
			US 2001-951265	A2 20010911

EP 2001-973722	A3 20010911
EP 2005-17665	A3 20010911
JP 2002-526851	A3 20010911
US 2002-116117	A 20020405
US 2002-284017	A1 20021030
WO 2003-US10463	W 20030404
US 2004-886950	A1 20040708
IN 2004-KN1494	A3 20041006

OTHER SOURCE(S) : MARPAT 138:153534
GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. of formulas I and II are provided [for I: Z = O, S, (un)substituted NH; Y = certain OH derivs., CHO, esters and amides of CO2H, certain NH2 derivs.; R1-R4 = H, halo, cyano, NO2, OH or derivs., NH2 or derivs., (un)substituted amidinyl, guanidinyl, alk(en/yn)yl, aryl, heterocycl1, CHO, CO2H and esters and amides; R5-R8 = H, halo, NO2, OH or derivs., NH2 or derivs., SH or derivs., cyano, etc.; R9 = H, OH, (un)substituted alkoxy or aryloxy, NH2 or derivs., (un)substituted alkyl or aryl, CHO, alkanoyl, aroyl; for II: A, B, D, E = C or N, with at least one being N; Y = H, OH or derivs., SH or derivs., NH2 or derivs., cyano, various acyl groups, (un)substituted alk(en/yn)yl, aralkyl, heterocycloalkyl, aryl, etc.; R1-R8 = H, halo, NO2, cyano, OH or derivs., NH2 or derivs., acyl, SH or derivs., etc.; R9 = H, OH, (un)substituted alkoxy, aryloxy, NH2 or derivs., aryl, CHO, alkanoyl, aroyl]. Also provided are pharmaceutical formulations including the compds. or their pharmaceutically acceptable salts and pharmaceutically acceptable carrier, which may be prepared by mixing the compds. or salts with a carrier and water. A disclosed method of treating a patient includes administering a pharmaceutical formulation according to the invention to a patient. Claims include tautomers of the compds., pharmaceutically acceptable salts, and pharmaceutically acceptable salts of the tautomers. I and II are inhibitors of receptor tyrosine kinases, and particularly of vascular endothelial growth factor receptor (VEGFR) tyrosine kinase. As such, they are inhibitors of angiogenesis, and thereby act as anticancer agents. Approx 270 invention compds. are listed, with detailed preps. given for about 50 compds. Several general preparatory methods are discussed in detail. For instance, cyclocondensation of Et 2-(benzimidazol-2-yl)acetate with the corresponding ortho-amino nitrile (preps. given), carried out in refluxing ClCH2CH2Cl in the presence of SnCl4, gave the invention quinolinone III. Many compds. I and II had in vitro IC50 values of less than 10 μ M with respect to flt-1 (VEGFR1), KDR (VEGFR2) and bFGF kinases (recombinant, expressed in Sf9 insect cells).

OS.CITING REF COUNT: 12 THERE ARE 12 CAPLUS RECORDS THAT CITE THIS RECORD (12 CITINGS)

L7 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2009 ACS on STN
 ACCESSION NUMBER: 2002:220574 CAPLUS
 DOCUMENT NUMBER: 136:263158
 TITLE: Benzimidazolyl-substituted quinolinone derivatives and analogs, with inhibitory action against vascular endothelial growth factor receptor tyrosine kinase, and useful as anticancer agents
 INVENTOR(S): Renhowe, Paul; Pecchi, Sabina; Machajewski, Tim; Shafer, Cynthia; Taylor, Clarke; McCrea, Bill; McBride, Chris; Jazan, Elisa; Wernette-Hammond, Mary-Ellen; Harris, Alex

PATENT ASSIGNEE(S): Chiron Corporation, USA
 SOURCE: PCT Int. Appl., 207 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 2
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002022598	A1	20020321	WO 2001-US42131	20010911 <--
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2421120	A1	20020321	CA 2001-2421120	20010911 <--
CA 2421120	C	20080715		
AU 2001093275	A	20020326	AU 2001-93275	20010911 <--
EP 1317442	A1	20030611	EP 2001-973722	20010911 <--
EP 1317442	B1	20051116		
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HU 2003001045	A2	20031229	HU 2003-1045	20010911 <--
BR 2001013757	A	20040302	BR 2001-13757	20010911 <--
JP 2004509112	T	20040325	JP 2002-526851	20010911 <--
JP 4361727	B2	20091111		
NZ 524717	A	20040924	NZ 2001-524717	20010911 <--
AU 2001293275	B2	20050414	AU 2001-293275	20010911 <--
AT 309996	T	20051215	AT 2001-973722	20010911 <--
ES 2250480	T3	20060416	ES 2001-973722	20010911 <--
EP 1650203	A1	20060426	EP 2005-17665	20010911 <--
EP 1650203	B1	20080220		
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AP 1666	A	20061031	AP 2003-2781	20010911 <--
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SG 129306	A1	20070226	SG 2005-1676	20010911 <--
EP 1849782	A1	20071031	EP 2007-11978	20010911 <--
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CN 100351249	C	20071128	CN 2001-815371	20010911 <--
AT 386736	T	20080315	AT 2005-17665	20010911 <--
ES 2302106	T3	20080701	ES 2005-17665	20010911 <--
ZA 2003001578	A	20040826	ZA 2003-1578	20030226 <--
IN 2003KN00244	A	20050311	IN 2003-KN244	20030226 <--
MX 2003002032	A	20030724	MX 2003-2032	20030307 <--
NO 2003001097	A	20030325	NO 2003-1097	20030310 <--
NO 324155	B1	20070903		
US 20040006101	A1	20040108	US 2003-387355	20030312 <--
US 6762194	B2	20040713		
BG 107709	A	20040130	BG 2003-107709	20030408 <--
HK 1053644	A1	20060504	HK 2003-104217	20030612 <--
US 20050054672	A1	20050310	US 2004-886950	20040708
US 7598268	B2	20091006		
HK 1064368	A1	20080926	HK 2004-106977	20040914
US 20050209456	A1	20050922	US 2005-92137	20050329
US 7335774	B2	20080226		

AU 2005202068	A1	20050602	AU 2005-202068	20050513
AU 2005202068	B2	20070809		
KR 2006036494	A	20060428	KR 2006-707122	20060413
KR 765841	B1	20071010	KR 2006-717401	20060828
JP 2007191486	A	20070802	JP 2007-62683	20070312
NO 2007001888	A	20030325	NO 2007-1888	20070412
IN 2008KN01705	A	20081226	IN 2008-KN1705	20080428
PRIORITY APPLN. INFO.:				
			US 2000-232159P	P 20000911
			AU 2001-293275	A3 20010511
			EP 2001-973722	A3 20010911
			EP 2005-17665	A3 20010911
			JP 2002-526851	A3 20010911
			US 2001-951265	A1 20010911
			WO 2001-US42131	W 20010911
			US 2002-284017	A1 20021030
			IN 2003-KN244	A3 20030226
			KR 2003-703558	A3 20030311
			US 2004-886950	A1 20040708

OTHER SOURCE(S): MARPAT 136:263158
GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. of formulas I and II are provided [for I: Z = O, S, (un)substituted NH; Y = certain OH derivs., CHO, esters and amides of CO2H, certain NH2 derivs.; R1-R4 = H, halo, cyano, NO2, OH or derivs., NH2 or derivs., (un)substituted amidinyl, guanidinyl, alk(en/yn)yl, aryl, heterocyclyl, CHO, CO2H and esters and amides; R5-R8 = H, halo, NO2, OH or derivs., NH2 or derivs., SH or derivs., cyano, etc.; R9 = H, OH, (un)substituted alkoxy or aryloxy, NH2 or derivs., (un)substituted alkyl or aryl, CHO, alkanoyl, aroyl; for II: A, B, D, E = C or N, with at least one being N; Y = H, OH or derivs., SH or derivs., NH2 or derivs., cyano, various acyl groups, (un)substituted alk(en/yn)yl, aralkyl, heterocycloalkyl, aryl, etc.; R1-R8 = H, halo, cyano, OH or derivs., NH2 or derivs., acyl, SH or derivs., etc.; R9 = H, OH, (un)substituted alkoxy, aryloxy, NH2 or derivs., aryl, CHO, alkanoyl, aroyl]. Also provided are pharmaceutical formulations including the compds. or their pharmaceutically acceptable salts and a pharmaceutically acceptable carrier, which may be prepared by mixing the compds. or salts with a carrier and water. A disclosed method of treating a patient includes administering a pharmaceutical formulation according to the invention to a patient. Claims include tautomers of the compds., pharmaceutically acceptable salts, and pharmaceutically acceptable salts of the tautomers. I and II are inhibitors of receptor tyrosine kinases, and particularly of vascular endothelial growth factor receptor (VEGFR) tyrosine kinase. As such, they are inhibitors of angiogenesis, and thereby act as anticancer agents. Approx 270 invention compds. are listed, with detailed preps. given for about 50 compds. Several general preparatory methods are discussed in detail. For instance, cyclocondensation of Et 2-(benzimidazol-2-yl)acetate with the corresponding ortho-amino nitrile (preps. given), carried out in refluxing ClCH2CH2Cl in the presence of SnCl4, gave the invention quinolinone III. Many compds. I and II had in vitro IC50 values of less than 10 μ M with respect to flt-1 (VEGFR1), KDR (VEGFR2) and bFGF kinases (recombinant, expressed in Sf9 insect cells).

OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS RECORD (10 CITINGS)

REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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FILE 'REGISTRY' ENTERED AT 13:59:00 ON 30 NOV 2009

L1 STRUCTURE UPLOADED
L2 1 S L1
L3 34 S L1 FULL
L4 1 S L1 SSS

FILE 'CAPLUS' ENTERED AT 14:00:29 ON 30 NOV 2009

L5 70 S L3
L6 54 S L5 AND (?CANCER? OR ?TUMOR? OR ?TUMOUR? OR ?NEOPLASM?)
L7 5 S L6 AND AD<20031107

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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
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CA SUBSCRIBER PRICE	-4.10	-4.10
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COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
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FULL ESTIMATED COST	5.25	231.13
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S L9
L10 112 L9

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2 FILES SEARCHED...
L12 5 L11 AND PD<20031107

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L12 ANSWER 1 OF 5 EMBASE COPYRIGHT (c) 2009 Elsevier B.V. All rights reserved on STN

ACCESSION NUMBER: 2003481481 EMBASE
TITLE: The impact of anti-angiogenic agents on cancer therapy.
AUTHOR: Marme, Dieter (correspondence)
CORPORATE SOURCE: Tumor Biology Center, Institute of Molecular Oncology, Breisacherstrasse 117, 79106 Freiburg, Germany. marme@tumor.bio.uni-freiburg.de
SOURCE: Journal of Cancer Research and Clinical Oncology, (Nov 2003) Vol. 129, No. 11, pp. 607-620.
Refs: 89
ISSN: 0171-5216 CODEN: JCROD7
COUNTRY: Germany
DOCUMENT TYPE: Journal; General Review; (Review)
FILE SEGMENT: 016 Cancer
030 Clinical and Experimental Pharmacology
037 Drug Literature Index
038 Adverse Reactions Titles
LANGUAGE: English
ENTRY DATE: Entered STN: 29 Dec 2003
Last Updated on STN: 29 Dec 2003

L12 ANSWER 2 OF 5 EMBASE COPYRIGHT (c) 2009 Elsevier B.V. All rights reserved on STN

ACCESSION NUMBER: 2003373828 EMBASE
TITLE: Anti-cancer drug discovery and development summit.
AUTHOR: Blakey, David C. (correspondence)

CORPORATE SOURCE: AstraZeneca, Alderley Park, Macclesfield, Cheshire SK10 4TF, United Kingdom. david.blakey@astrazeneca.com
SOURCE: Expert Opinion on Investigational Drugs, (1 Sep 2003) Vol. 12, No. 9, pp. 1577-1582.
Refs: 15
ISSN: 1354-3784 CODEN: EOIDER
COUNTRY: United Kingdom
DOCUMENT TYPE: Journal; Conference Article; (Conference paper)
FILE SEGMENT: 016 Cancer
030 Clinical and Experimental Pharmacology
037 Drug Literature Index
038 Adverse Reactions Titles
LANGUAGE: English
SUMMARY LANGUAGE: English
ENTRY DATE: Entered STN: 2 Oct 2003
Last Updated on STN: 2 Oct 2003
AB The 5th Annual Anti-Cancer Drug Discovery and Development Summit brought together an international group of academic and industry scientists to discuss recent therapeutic developments in the field of oncology. The focus of the meeting was novel targeted approaches, i.e., those agents directed against targets that are overexpressed or overactive in tumour cells. It was acknowledged that cytotoxic agents will continue to play a key role in the treatment of cancer and new developments in this area were also discussed. With over 400 anticancer drugs in clinical development and a number of recent registrations, there is great optimism that significant therapeutic advances can be made.
L12 ANSWER 3 OF 5 EMBASE COPYRIGHT (c) 2009 Elsevier B.V. All rights reserved on STN
ACCESSION NUMBER: 2003363876 EMBASE
TITLE: American Association for Cancer Research - 9th Annual Meeting: Investigating drugs: 11-14 July 2003, Washington, DC, USA.
AUTHOR: Mackay, Janie (correspondence); Williams, Laura
CORPORATE SOURCE: Thomson Current Drugs, Middlesex House, 34-42 Cleveland Street, London W1T 4JE, United Kingdom. laura.williams@current-drugs.com
SOURCE: IDrugs, (1 Aug 2003) Vol. 6, No. 8, pp. 736-738.
ISSN: 1369-7056 CODEN: IDRUFN
COUNTRY: United Kingdom
DOCUMENT TYPE: Journal; Conference Article; (Conference paper)
FILE SEGMENT: 016 Cancer
030 Clinical and Experimental Pharmacology
036 Health Policy, Economics and Management
037 Drug Literature Index
038 Adverse Reactions Titles
052 Toxicology
LANGUAGE: English
ENTRY DATE: Entered STN: 25 Sep 2003
Last Updated on STN: 25 Sep 2003
L12 ANSWER 4 OF 5 EMBASE COPYRIGHT (c) 2009 Elsevier B.V. All rights reserved on STN
ACCESSION NUMBER: 2003276961 EMBASE
TITLE: Kinases - SMi Conference 9-10 April 2003, London, UK.
AUTHOR: Harrison, Ruth (correspondence)
CORPORATE SOURCE: Thomson Current Drugs, Middlesex House, 34-42 Cleveland Street, London W1T 4LB, United Kingdom. ruth.harrison@current-drugs.com
SOURCE: IDrugs, (1 Jun 2003) Vol. 6, No. 6, pp. 560-562.
ISSN: 1369-7056 CODEN: IDRUFN

COUNTRY: United Kingdom
 DOCUMENT TYPE: Journal; Conference Article; (Conference paper)
 FILE SEGMENT: 029 Clinical and Experimental Biochemistry
 030 Clinical and Experimental Pharmacology
 031 Arthritis and Rheumatism
 037 Drug Literature Index
 LANGUAGE: English
 SUMMARY LANGUAGE: English
 ENTRY DATE: Entered STN: 24 Jul 2003
 Last Updated on STN: 24 Jul 2003
 AB Dr. Moss briefly summed up the conference by describing the growth in the development of kinase research over the years and the commitment being invested by companies aiming to find effective screening strategies. He closed the day by remarking on the new challenge for researchers of turning the concepts discussed into successful drugs.
 L12 ANSWER 5 OF 5 BIOSIS COPYRIGHT (c) 2009 The Thomson Corporation on STN
 ACCESSION NUMBER: 2003:501918 BIOSIS
 DOCUMENT NUMBER: PREV200300498316
 TITLE: Preclinical pharmacokinetics and metabolism of CHIR258, a potent tyrosine kinase inhibitor.
 AUTHOR(S): Vora, Jayesh [Reprint Author]; Haroldsen, Peter [Reprint Author]; Renhove, Paul [Reprint Author]; Heise, Carla [Reprint Author]; Steigerwalt, Ronald [Reprint Author]; Todd, Marque [Reprint Author]; Harris, Alex [Reprint Author]; Samara, Emil [Reprint Author]
 CORPORATE SOURCE: Chiron Corporation, Emeryville, CA, USA
 SOURCE: Proceedings of the American Association for Cancer Research Annual Meeting, (July 2003) Vol. 44, pp. 753. print.
 Meeting Info.: 94th Annual Meeting of the American Association for Cancer Research. Washington, DC, USA. July 11-14, 2003.
 ISSN: 0197-016X.
 DOCUMENT TYPE: Conference; (Meeting)
 Conference; Abstract; (Meeting Abstract)
 LANGUAGE: English
 ENTRY DATE: Entered STN: 29 Oct 2003
 Last Updated on STN: 29 Oct 2003

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FULL ESTIMATED COST	ENTRY	SESSION	
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CA SUBSCRIBER PRICE	ENTRY	SESSION	
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L13 1 CHIR 258/CN

=> d 113

L13 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2009 ACS on STN
RN 692737-80-7 REGISTRY
ED Entered STN: 14 Jun 2004
CN Propanoic acid, 2-hydroxy-, compd. with
4-amino-5-fluoro-3-[6-(4-methyl-1-piperazinyl)-1H-benzimidazol-2-yl]-2(1H)-
quinolinone (1:1) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Propanoic acid, 2-hydroxy-, compd. with
4-amino-5-fluoro-3-[5-(4-methyl-1-piperazinyl)-1H-benzimidazol-2-yl]-2(1H)-
quinolinone (1:1) (9CI)

OTHER NAMES:

CN CHIR 258

CN Dovitinib lactate

CN TKI 258

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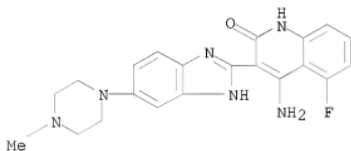
SR CA

LC STN Files: CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL

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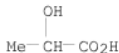


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PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

61 REFERENCES IN FILE CA (1907 TO DATE)
 2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
 63 REFERENCES IN FILE CAPLUS (1907 TO DATE)

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	ENTRY	SESSION	
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FILE COVERS 1907 - 30 Nov 2009 VOL 151 ISS 23
 FILE LAST UPDATED: 29 Nov 2009 (20091129/ED)
 REVISED CLASS FIELDS ('NCL') LAST RELOADED: Aug 2009
 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

CAPplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

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 L14 63 L13

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L16 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN
 ACCESSION NUMBER: 20051242789 CAPLUS
 DOCUMENT NUMBER: 143:477969
 TITLE: Preparation of benzimidazole quinolinones for
 inhibiting FGFR3 and treating multiple myeloma
 INVENTOR(S): Cai, Shaopei; Chou, Joyce; Harwood, Eric; Heise, Carla
 C.; Machajewski, Timothy D.; Ryckman, David; Shang,
 Xiao; Wiesmann, Marion; Zhu, Shuguang
 PATENT ASSIGNEE(S): Chiron Corporation, USA
 SOURCE: U.S. Pat. Appl. Publ., 239 pp., Cont.-in-part of U.S.
 Ser. No. 644,055.
 CODEN: USXXCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 7
 PATENT INFORMATION:

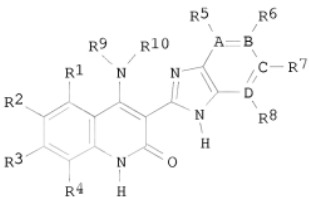
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US 20050261307	A1	20051124	US 2004-983174	20041105
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US 7470709	B2	20081230		
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ZA 2006003598	A	20080430	ZA 2006-3598	20060505
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US 20090181979	A1	20090716	US 2009-398130	20090304
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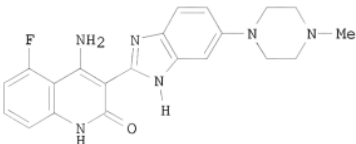
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 143:477969

GI



I



II

AB The title compds. I [A, B, C, and D = C, N; R1-R3 = H, halo, CN, NO2, etc.; R4 = H, alkyl; R5-R8 = H, halo, CN, NO2, etc.; R9 = H, (un)substituted alkyl, aryl, etc.; R10 = H], useful for inhibiting fibroblast growth factor receptor 3 or treating a biol. condition mediated by fibroblast growth factor receptor 3, were prepared E.g., a multi-step synthesis of 4-amino-5-fluoro-3-[6-(4-methylpiperazin-1-yl)-1H-benzimidazol-2-yl]-1H-quinolin-2-one (II), starting from 5-chloro-2-nitroaniline and 1-methylpiperazine, was given. The majority of the exemplary compds. I displayed an IC50 of less than 10 μ M with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, Cdk4, MEK1, NEK-2, CHK2, CK1 α , Raf, Fyn, Lck, Rsk2, PAR-1, c-Kit, c-ABL, p60 src , FGFR3, FLT-3, PDGFR α , and PDGFR β . In addition, many of the exemplary compds. exhibited IC50 values in the nM range and show potent activity with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, FGFR3, c-Kit, c-ABL, FLT-3, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, MEK1, CHK2, Fyn, Lck, Rsk2, PAR-1, PDGFR α , and PDGFR β with IC50 values of less than 1 μ M. The mentioned above compound II was tested in various tests and showed significant antiproliferative activity. II inhibited FGFR3 receptor phosphorylation and ERK phosphorylation in multiple myeloma cell lines with activating FGFR3 mutations.

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

L16 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2005:1223876 CAPLUS

DOCUMENT NUMBER: 143:477966

TITLE: Preparation of benzimidazole quinolinones for inhibiting a checkpoint kinase 1 and their use in combination therapy for cancer

INVENTOR(S): Gesner, Thomas G.; Barsanti, Paul A.; Harrison, Stephen D.; Ni, Zhi-Jie; Brammeier, Nathan M.; Zhou, Yasheen; Le, Vincent P.

PATENT ASSIGNEE(S): Chiron Corporation, USA

SOURCE: U.S. Pat. Appl. Publ., 249 pp., Cont.-in-part of U.S. Ser. No. 644,055.

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 7

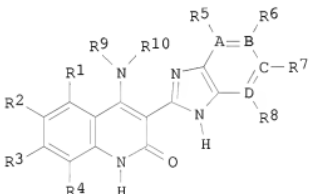
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US 20050256157	A1	20051117	US 2005-41191	20050121
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US 7470709	B2	20081230		
CN 1692112	A	20051102	CN 2003-824565	20030819 <--
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		US 2003-480448P	P	20030701
		US 2003-644055	A2	20030819
		US 2004-538984P	P	20040123

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): CASREACT 143:477966; MARPAT 143:477966

GI



AB The title compds. [I; A, B, C, D = C, N; R1 = H, halo, CN, NO₂, etc.; R2, R3 = H, halo, NO₂, CN, etc.; R4 = H, (un)substituted alkyl; R5, R8 = H, (un)substituted alkyl, alkenyl, heterocycl; or R5 may be absent if A = N; or R8 may be absent if D = N; R6, R7 = H, halo, NO₂, CN, etc.; R9 = H, (un)substituted alkyl, aryl, etc.; R10 = H; or R9 and R10 join together to form one or more rings, each having 5-7 members], useful for inhibiting checkpoint kinase 1, inducing cell cycle progression, and increasing apoptosis in cells, were prepared. E.g., a multi-step synthesis of 4-amino-3-(benzimidazol-2-yl)-6-(4-methylpiperazinyl)hydroquinolin-2-one, was given. The compds. I were tested against various kinases. Two of the prepared compds. I, 4-[(3S)-1-azabicyclo[2.2.2]oct-3-ylamino]-3-(1H-

benzimidazol-2-yl)-6-chloroquinolin-2-(1H)-one and 6-chloro-3-[5-(4-methylpiperazin-1-yl)-1H-benzimidazol-2-yl]-4-[(piperidin-2-ylmethyl)amino]quinolin-2(1H)-one, were found to be potent inhibitors of CHK1 with IC50 of 0.32 nM and 0.63 nM, resp. The majority of the exemplary compds. I displayed an IC50 of less than 10 μ M with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, Cdk4, MEK1, NEK-2, CHK2, CKit, Raf, Fyn, Lck, Rsk2, PAR-1, c-Kit, c-ABL, p60src, FGFR3, FLT-3, PDGFR α , and PDGFR β . In addition, many of the exemplary compds. exhibited IC50 values in the nM range and show potent activity with respect to VEGFR1, VEGFR2, VEGFR3, FGFR1, FGFR3, c-Kit, c-ABL, FLT-3, CHK1, Cdc2, GSK-3, NEK-2, Cdk2, MEK1, CHK2, Fyn, Lck, Rsk2, PAR-1, PDGFR α , and PDGFR β with IC50 values of less than 1 μ M. The compds. I may be used to prepare pharmaceutical compns. and may be used in conjunction with DNA damaging agents.

OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

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L16 2 S L15 AND AD<20031107

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